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Description

The present invention relates to a method for forming an article by exposing a polymer to radiation having a wavelength of about 10^{-3} to about 265 nm to cure or cross-link the polymer, and to articles containing radiation cross-linked polymers.

Many manufacturing processes employ radiation to cure or cross-link polymers. Exemplary examples of radiation are ultraviolet (UV) light having a wavelength of about 10⁻² to about 10 nm, and electron beams having a wavelength of about 10⁻³ to about 10⁻¹ nm. The polymers can cross-link with themselves to increase their molecular weight. Typically, the resulting polymer exhibits new, and often more desirable, properties. Alternatively, the polymer can cross-link with or be covalently immobilized onto a substrate. This is desirable for those applications where a tenacious coating or bond is desired.

There are advantages of using radiation to cross-link polymers. For example, products for use in certain fields, e.g., health care, must be sterilized. One method of sterilizing articles involves exposing them to UV light. Products made with non-UV light curable polymers are first manufactured and then sterilized. This two-step procedure adds to the cost of the finished product. Although some polymers are radiation curable, most polymers are not.

Accordingly, there is a need for radiation-curable polymers that can be employed in radiation-curable methods for making articles, e.g., articles suitable for use in fields requiring sterilized products.

We have discovered that a polymer comprising a backbone and a plurality of pendant functional groups having the formula

in which R₁ is a divalent radical and X is organoacyl or cyano, is radiation curable and may be employed in the manufacture of articles by methods involving exposure to radiation to accomplish curing of the polymer to meet the foregoing needs.

Polymers having the foregoing structure and in which R₁ is at least 3 atoms in length are known from EP-A-0 241 127 that discloses their use as a binder for woven and non-woven fibre assemblies. However it has not hitherto been known or suspected that such polymers are curable by UV or shorter wave length radiation.

The invention accordingly provides a method as defined in the claims hereof. In this method radiation having a wavelength within the range about 10⁻³ to about 265 nm is utilized to accomplish curing of the polymer. Generally UV light having a wavelength in the range 10 to 265 nm is used.

The radiation-curable polymer can be made by reacting a plurality of monomers with at least one polymerizable functional monomer having the formula

$$R_9 - CH = C - R_1 - C - CH_2 - X$$

wherein X is as defined above and R₂ and R₁₀ are each independently selected from the group consisting of hydrogen, halo, thio, and monovalent organic radicals.

Articles produced by the method of the instant invention contain polymers that are cross-linked to themselves and/or covalently bonded to a substrate. In addition, in one embodiment of the invention when the articles are exposed to UV light during their manufacture, the articles are sterilized and suitable for use in applications requiring sterilized goods.

The present invention is directed to (a) a method for forming radiation-cured articles and (b) the articles produced thereby. More particularly, the method of the instant invention comprises the steps of (a) contacting a substrate with a polymer to form a substrate/polymer combination, and (b) curing or cross-linking the polymer by exposing the substrate/polymer combination to radiation having a wavelength within the said range and capable of cross-linking the polymer. The polymer comprises a backbone and a plurality of pendant functional groups. In one version, the backbone of the polymer is formed by combining a

plurality of olefinically unsaturated monomers. Exemplary monomers are olefinically unsaturated carboxylic acid ester monomers and salts thereof. Alternatively, the backbone of the polymer is formed by other polymerization reactions, e.g., condensation reactions.

The pendant functional groups have the formula

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$$-R_1 - C - CH_2X \qquad (1)$$

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wherein R₁ is a divalent radical, and X is selected from the group consisting of organoacyl and cyano. R₁ can be or contains heteroatoms, such as oxygen, sulfur, phosphorus, and nitrogen. The heteroatoms are preferably selected from the group consisting of oxygen, sulfur, and nitrogen. In addition, R1 can contain functional groups such as carbonyls, carboxy-esters, thio, and amino substituents. Although R1 can also comprise aromatic, olefinic, or alkynyl unsaturation, R1 is preferably saturated. Preferably, R1 is a cyclic or acyclic divalent organic radical containing up to about 40 carbon atoms. Most preferably, Rt is acyclic containing up to about 20 atoms in length, with any and all side groups each being up to about 6 atoms in length. Because of commercial availability, X preferably is organoacyl. X generally has the formula

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$$\begin{array}{ccc}
0 \\
- C - R_2
\end{array} \tag{II}$$

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wherein R2 is selected from the group consisting of hydrogen and monovalent organic radicals. (As used throughout the specification and claims, the term "organic radical" refers to any group containing at least one carbon atom, e.g., aliphatic and aromatic radicals, whether containing only hydrogen and carbon (i.e., hydrocarbon radicals) or further containing heteroatoms such as oxygen, sulfur, and nitrogen, and/or an inorganic substituent such as chlorine, bromine, and iodine. Accordingly, organic radicals include, for example, substituted and unsubstituted alkyl, aryl, arylalkyl, alkylaryl, alkyloxy, aryloxy, arylalkyloxy, alkenyl, alkenyloxy, alkyni, alkynyloxy, and arylalkenyl radicals, and heteroatom-containing hydrocarbyl radicals. The heteroatoms are preferably selected from oxygen, sulfur, and nitrogen atoms.) Typically, R2 contains up to about 10 atoms in addition to any hydrogen atoms present in the monovalent organic radical. Preferably, R2 is H or an alkyl group containing up to about 6 carbon atoms. Methyl is most preferred for R2.

An exemplary pendant functional group has the formula

$$\frac{0}{1} - \frac{0}{1} - \frac{0}{1} - \frac{0}{1} - \frac{0}{1} - \frac{0}{1} - \frac{0}{1} = \frac{0}{1}$$
 (III)

wherein R₃ is a divalent organic radical at least 2 atoms in length, Y and Z are each independently selected from the group consisting of O, S, and NR4, with R4 being selected from the group consisting of H and hydrocarbyl radicals containing up to about 6 carbon atoms. Preferably, R4 is H or an alkyl group. Oxygen is preferred for Y and Z. Typically, R3 contains up to about 40 atoms, but usually contains no more than about 20 atoms. R₃ can be cyclic or acyclic or can contain both cyclic and acyclic moleties. Exemplary cyclic R₃ groups include cycloalkylenes and phenylene. R₃ is preferably acyclic and is selected from the groups consisting of substituted and unsubstituted alkylenes, polyoxyalkylenes, polythioalkylenes, and polyaminoalkylenes. Unsubstituted alkylenes are most preferred for R₃. The substituted and unsubstituted polythio-, polyoxy-, and polyaminoalkylenes can be formed by the well known condensation of alkylene oxides, alkylene amines, glycols, diamines, and dithiols. For example, polyoxyalkylenes can be formed as follows:

$$n(R_5 - CH_2) \rightarrow HO(CH_2R_5 CH_2 - O)_nH$$
 (IV)

wherein R₅ is H or a monovalent organic radical and the product of formula IV represents HO(CHR₅ - CH₂ - O)_xH, HO(CH₂ - CHR₅ - O)_nH, and mixtures thereof R₅ contains up to about 40 carbon atoms, but generally contains less than about 20 carbon atoms. Preferably, R₅ is H or an alkyl group containing up to about 10 carbon atoms.

Other exemplary R₃ radicals are:

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wherein m is 0 or 1, n is 0 or 1, and R_{ϵ} is a divalent organic radical at least 1 atom in length, and R_7 is a divalent organic radical at least 1 atom in length. Typically R_{ϵ} and R_7 are alkylene groups containing up to about 3 carbon atoms in length.

The pendant functional groups are attached to the polymer backbone either as part of a polymerizable olefinically unsaturated monomer or by substitution onto a polymer by any suitable addition reaction. Typical addition reactions are schematically represented as follows:

Polymer backbone
$$(-C-C1)_p + (H-O-R_8-C-CH_2-X)_p \rightarrow$$

Polymer backbone - (Cl)_p + (H - O - R₈ -
$$\ddot{c}$$
 - CH₂ - x)_p \longrightarrow

where p is an integer and - O - R_8 is R_1 in formula I, supra.

When the pendant functional groups are incorporated into the polymer backbone as part of a polymerizable monomer, the polymerizable monomer can have the formula

$$R_9 - CH = C - R_1 - C - CH_2 - X$$
 (VIII)

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wherein X is as defined in formula I, supra, R_9 and R_{10} are each independently selected from the group consisting of hydrogen, halo, thio, and monovalent organic radicals. Generally, R_9 and R_{10} contain up to about 10 atoms other than hydrogen. Preferably, R_9 and R_{10} are H and alkyl radicals having up to about 10 carbon atoms. An exemplary polymerizable monomer has the formula

$$R_9 - CH = C - C - Y - R_3 - Z - C - CH_2 - C - R_2$$
 (IX)

wherein Y, Z, R₂, R₃, R₉, and R₁₀ are as defined above. A preferred class of polymerizable monomers due to their commercial availability, can be represented by the following formula:

$$CH_{2} = \overset{R}{C} - \overset{O}{C} - O - R_{3} - O - \overset{O}{C} - CH_{2} - \overset{O}{C} - R_{2}$$
 (X)

wherein R_2 is an alkyl group containing up to about 8 carbon atoms, R_3 is an alkylene group containing up to about 10 carbon atoms, and R_{10} is hydrogen or an alkyl group containing up to about 12 carbon atoms. Exemplary monomers of formula X include:

$$CH_{2} = \overset{CH_{3}}{C} - \overset{O}{C} - O - CH_{2}CH_{2} - O - \overset{O}{C} - CH_{2} - \overset{O}{C} - CH_{3}$$
 (XI)

$$\frac{CH_{3}}{CH_{2}} = \overset{C}{C} - \overset{C}{C} - O - CH_{2}CH_{2}CH_{2} - O - \overset{O}{C} - CH_{2} - \overset{O}{C} - CH_{3} \qquad (XII)$$

$$CH_{2} = \overset{CH_{3}}{c} - \overset{O}{c} - O - CH_{2} \overset{CH_{2}}{CH_{2}} - O - \overset{O}{c} - CH_{2} - \overset{O}{c} - CH_{3}$$
(XIII)

The polymer contains a sufficient amount of one or more of the pendant functional groups of formula I to enable the polymer to be radiation cross-linkable. Generally, the polymer contains at least about 0.5 weight percent of the pendant functional groups based on the total polymer content. More typically, the polymer contains at least about one weight percent of the pendant functional groups of formula I, supra. Usually, a pendent functional group concentration in excess of about 20 weight percent does not produce significantly greater technical benefits. Therefore, the pendant or functional groups are normally present in the polymer in a concentration of about 0.5 to about 20 weight percent and, more commonly, in the range of about 0.5 to about 10 weight percent based on the total polymer content. Pendant functional groups containing different R₁ and X groups can be blended in the same polymer. Alternatively, polymers containing different R₁ and X groups can be blended in the same solution or colloid.

Radiation used for cross-linking the polymer has a wavelength in the range of about 10^{-3} to about 265 nm. This range includes UV light having a wavelength of about 10 to about 265 nm, X-rays having a wavelength of about 10^{-2} to about 10 nm, and electron beams having a wavelength of about 10^{-3} to about

10 nm. UV light and electron beams are preferred radiation sources because they are more commonly used. In particular, UV is extensively used in industry. Since the energy of the UV light increases as the UV wavelength decreases, it is preferred that the UV light have a wavelength in the range of about 10 to about 250, and most preferably in the range of about 10 to about 225 nm. However, it is believed that satisfactory cross-linking is obtained with UV light having a wavelength in the range of about 225 to about 265 nm.

The polymer can be applied to the substrate as part of an aqueous colloid or a solvent-containing solution. Alternatively, the polymer can be applied "neat", that is, substantially by itself.

Aqueous colloids and solvent-containing solutions of the polymers can be prepared by procedures known in the art. For instance, aqueous polymer colloids can be prepared by gradually adding each monomer simultaneously to an aqueous reaction medium at rates proportionate to the respective percentage of each monomer in the finished polymer. Polymerization is initiated and continued by providing a suitable polymerization catalyst in the aqueous reaction medium. Illustrative polymerization catalysts are free radical initiators and redox systems such as hydrogen peroxide, potassium and ammonium peroxydisulfate, dibenzoyl peroxide, hydrogen peroxide, lauryl peroxide, di-tertiary-butyl peroxide, and azobisisobutyronitrile. These catalysts can be employed either alone or together with one or more reducing components such as sodium bisulfate, sodium metabisulfate, glucose, ascorbic acid, and erythorbic acid. The polymerization reaction is continued with agitation at a temperature sufficient to maintain an adequate reaction rate until all added monomers are consumed. Monomer addition is usually continued until the latex (colloid) reaches a polymer concentration of about 10 to about 65 weight percent. Typically, latexes have a solids content of about 40 to about 60 weight percent.

Physical stability of the colloid is achieved by providing one or more surfactants (emulsifiers) in the aqueous reaction medium. Non-ionic, anionic, and/or amphoteric surfactants can be employed. Exemplary non-ionic surfactants are alkylpolyglycol ethers such as ethoxylation products of lauryl, cleyl, and stearyl alcohols or mixtures of such alcohols such as coconut fatty alcohol; and alkylphenol polyglycol ethers such as ethoxylation products of octyl- or nonylphenol, diisopropylphenol, triisopropylphenyl, and di- or tritertiarybutylphenol. Examples of anionic surfactants are alkali metal and ammonium salts of alkyl, aryl, and alkylaryl sulfonates, sulfates, phosphates, phosphonates, and carboxylates. Specific examples of anionic surfactants include, but are not limited to, sodium lauryl sulfate, sodium octylphenyl glycolether sulfate, sodium dodecylbenzene sulfonate, sodium lauryldiglycol sulfate, ammonium tritertiarybutylphenol, penta- and octa-glycol sulfates, and polyacrylic acid. Betaines and aminosulfonates are examples of amphoteric surfactants. Numerous other examples of suitable anionic, non-ionic, and amphoteric surfactants are disclosed in U.S. Patents 2,600,831; 2,271,622; 2,271,623; 2,275,727; 2,787,604; 2,816,920; and 2,739,891, which references are incorporated herein by reference.

Colloidal stabilizing agents are optionally added to the aqueous polymer colloids either during or after the reaction period. Exemplary colloidal stabilizing agents include gum arabic, starch, alginates, and modified natural substances such as methyl-, ethyl-, hydroxyalkyl-, and carboxymethyl- cellulose, and synthetic substances such as polyvinyl alcohol, polyvinyl pyrrolidone, and mixtures thereof. Fillers and/or extenders, such as dispersible clays, and colors, such as pigments and dyes, can also be added to the aqueous colloid either during or after polymerization.

Another advantage of the present invention is that solutions and colloids, and particularly colloids in aqueous medium, of the polymers employed in the instant invention have a lower viscosity than ester polymers not containing the functional monomers useful in this invention. Thus, the latexes have viscosities of about 360 kg/m-h (100 centipoise)or less, often about 180kg/m-h (50 centipoise)or less, measured at 21 °C. at a polymer concentration of 40 weight percent or more and even at about 50 weight percent or more.

The low viscosity behavior of the concentrated latexes employed in the present invention is atypical, particularly for polymers having comparable molecular weights and for latexes of comparable molecular size. Generally, polymer molecular weight maximums are about 150,000 or less, and typically about 100,000 or less. The dispersed polymer particles can be of any size suitable for an intended use. However, since latex viscosity increases as particle size is reduced substantially below 120 nm, it is preferred that the polymer particle size be at least about 120 nm. Typically, the latexes have polymer particle sizes within the range of about 120 to about 300 nm as determined on the N-4 "Nanosizer" available from Cotter Electronics, Inc. of Hialeah, Florida.

Due to the lower viscosity of solutions and colloids containing the polymers used in the instant invention, the polymer content of both the aqueous colloids and solutions can be increased without exceeding permissible viscosity limits. Similarly, the loading of the colloids and solutions with fillers such as clays, pigments, and other extenders can be increased. For instance, aqueous dispersions and polymer solutions can contain more than about 2 percent, often more than 5 percent, and even more than 10 percent fillers, colorants, and/or extenders.

Solutions of the polymers can be prepared by polymerizing the selected monomers in solvents in which both the monomers and the polymers are soluble. Suitable solvents include aromatic solvents and alcohols. Xylene and toluene are exemplary aromatic solvents. An exemplary alcohol is butanol. Polymerization initiators and reducing components, when employed, should be soluble in the selected solvent or mixture of solvents. Suitable polymerization initiators soluble in the above organic solvents include dibenzoyl peroxide, lauryl peroxide, and azobisisobutyronitrile. Erythobic and ascorbic acids are exemplary reducing components that are soluble in polar organic solvents.

In addition to being UV light curable, polymers containing the pendant functional groups of formula I, supra, also improve one or more physical properties of substrates relative to a similar substrate containing a similar polymer absent such pendant functional groups. For example, the pendant functional groups produce significant improvements in substrate properties when employed with polymers which contain significant amounts of polymerized, olefinically, unsaturated mono- and/or polycarboxylic acid esters and/or their salts. These latter polymers usually contain at least about 10 weight percent, often at least about 20 weight percent, and typically at least about 30 weight percent of olefinically unsaturated, carboxylic acid ester monomers other than the functional monomers of formula VIII, supra. Generally, the polymers contain at least about 50 weight percent, and commonly at least about 80 weight percent, of such ester comonomers. Exemplary ester comonomers are esters of olefinically unsaturated mono- or dicarboxylic acids and/or their salts having up to about 10 carbon atoms, and hydroxy-, amino-, or thio-substituted or unsubstituted alcohols, amines, and thiols having from 1 to about 30 carbon atoms, usually 1 to about 20 carbon atoms, per molecule. Illustrative unsaturated carboxylic acids are acrylic, methacrylic, fumaric, maleic, and itaconic acid. Illustrative hydroxy-, amino-, and thio-substituted alcohols, amines, and thiols are glycerol, 1-hydroxy-5-thiododecane, and 2-amino-5-hydroxyhexane. Preferred ester comonomers, due primarily to cost and availability, are hydroxy-substituted and unsubstituted alcohol esters of acrylic and methacrylic acids such as butyl acrylate, 2-ethylhexyl acrylate, methyl methacrylate, and hydroxyethyl acrylate. Other desirable comonomers include acrylonitrile and acrylamide.

The functional monomers of formula VIII, supra, and the ester monomers can constitute the total polymer composition. Alternatively, the portion of the polymer molecule not accounted for by these two monomer classes can be any polymerizable, olefinically unsaturated monomer or combination of monomers. Illustrative of such other polymerizable monomers are vinyl esters of carboxylic acids, the acid moiety of which contains from 1 to about 20 carbon atoms (e.g., vinyl acetate, vinyl propionate, and vinyl isononoate); aromatic and aliphatic, alpha-beta-unsaturated hydrocarbons such as ethylene, propylene, styrene, and vinyl toluene; vinyl halides such as vinyl chloride and vinylidene chloride; olefinically unsaturated nitriles such as acrylonitrile; and olefinically unsaturated carboxylic acids having up to about 10 carbon atoms such as acrylic, methacrylic, crotonic, itaconic, and maleic acids. It has been found that minor amounts of olefinically unsaturated carboxylic acids and/or sulfoalkyl esters of such carboxylic acids and/or their salts significantly improve tensile strength and/or other physical properties of the finished textile material. Thus, it is preferred that the polymer contain at least about 0.1 weight percent, usually about 0.1 to about 10 weight percent, and typically about 0.1 to about 5 weight percent of (a) a polymerizable, olefinically unsaturated carboxylic acid having up to about 10 carbon atoms and/or (b) a sulfoalkyl ester of such acids such as sulfoethyl methacrylate, sulfoethyl itaconate, sulfomethyl malonate, and 2-acrylamido-2methylpropane sulfonic acid, and/or (c) salts of (a) and/or (b).

As noted above, substrates that are suitable for use in the present invention include textiles. Textile substrates include woven and non-woven fibers. For the purpose of this invention, the term "fibers" encompasses relatively short fibers as well as longer fibers often referred to as "filaments". Fibers contain either polar or non-polar functional groups. Exemplary polar functional groups contained in fibers are hydroxy, etheral, carbonyl, carboxylic acid (including carboxylic acid salts), carboxylic acid esters (including thio esters), amides, and amines. Essentially all natural fibers, e.g., virgin wool and reclaimed cellulosic fibers such as cotton, wood fiber, coconut fiber, jute, hemp, and protenaceous materials such as wool and other animal fur, include one or more polar functional groups. Exemplary synthetic fibers containing polar functional groups are polyesters, polyamides, and carboxylated styrene-butadiene polymers. Typical polyamides include nylon-6, nylon-66, an nylon-610. Typical polyesters include Dacron, Fortrel, and Kodel brand polyesters. Typical acrylic fibers include Acrilan, Orlon, and Creslen brand acrylic fibers. Exemplary modacrylic fibers include Verel and Dynel brand modacrylic fibers. Other useful fibers which are also polar include synthetic carbon, silicon, and magnesium silicate (e.g. asbestos) polymer fibers and metallic fibers such as aluminum, gold, and iron fibers. Exemplary non-polar functional groups containing fibers are polyolefin and styrene-butadiene.

Fibers containing polar functional groups are widely employed in manufacturing a vast variety of textile materials, including wovens, nonwovens, knits, threads, yarns, and ropes. Minor amounts of polymers

containing the pendant functional group of formula I, supra, are sufficient to enhance the physical properties of such articles, in particular tensile strength, abrasion resistance, scrub resistance, and/or shape retention, with little or no degradation of other desirable properties such as hand, flexibility, elongation, and physical and color stability. Polymer concentrations of at least about 5, generally at least about 8 weight percent based on the dry weight of the finished polymer-coated textile, are sufficient to obtain detectable physical property improvements in many textiles. Because of cost considerations, the polymer concentration rarely exeeds about 25 weight percent based on the dry weight of the finished polymer-coated textile. Typically, the polymer concentration is less than about 20 weight percent. Satisfactory results are achieved when the polymer concentration is within the range of about 10 to about 15 weight percent.

Although significantly greater improvements in physical properties are achieved by application of the polymers to polar group-containing fibers in contrast to relatively non-polar fibers, non-polar fibers can also be employed in the present invention. Furthermore, polar groups, such as carbonyl (e.g., keto) and hydroxy groups, can be introduced into polyclefins, styrene-butadiene polymers and other relatively non-polar fibers by known oxidation techniques.

The product property in which the most significant improvement results depends, to some extent, on the structure of the treated fiber assemblage. For instance, threads and ropes formed from relatively long, tightly wound or interlaced fibers and tightly woven textiles generally possess significant tensile strength in their native state. Accordingly, the percentage increase in tensile strength resulting from polymer treatment will be less, on a relative basis, than it is with other products such as loose-wovens, knits, and non-wovens which have a relatively tower tensile strength in their native form. However, significant improvements in abrasion resistance and scrub resistance are achieved in threads, ropes, and tightly woven textiles. Furthermore, in loose-woven textiles and knitted fabrics, significant improvements can be achieved for shape retention (including retention of the relative spacing of adjacent woven strands), abrasion resistance, and scrub resistance.

Significant advantages of the present invention are also obtained for non-woven fibers. Non-woven fibers depend primarily on the strength and persistence of the fiber-polymer bond for their physical properties and for the retention of such properties with use. Bonded non-woven fabrics, such as the textile articles of this invention, can be defined generally as assemblies of fibers held together in a random or oriented web or mat by a bonding agent. While many non-woven materials are manufactured from crimped fibers having lengths of about 13 to about 130mm (about 0.5 to about 5 inches), shorter or longer fibers can be employed. Exemplary utilities for non-woven textiles are hospital sheets, gowns, masks, as well as roadbed underlayment supports, diapers, roofing materials, napkins, coated fabrics, papers of all varieties, and tile backings (for ungrouted tile prior to installation). The physical properties of non-woven fibers range all the way from stiff, board-like homogeneous and composite paper products to soft drapeable textiles (e.g., drapes and clothing), and wipes.

Non-woven products can be generally divided into categories characterized as "flat" and "highloft" goods. Each category includes both disposable and durable products. Presently, the major end uses of disposable flat non-woven goods include diaper cover stock, surgical drapes, gowns, face masks, bandages, industrial work clothes, consumer and industrial wipes and towels such as paper towels, and feminine hygiene products. Current major uses of durable flat non-woven goods include apparel interlinings and interfacings, drapery and carpet backings, automotive components (such as components of composite landau automobile tops), carpet and rug backings, and construction materials, such as roadbed underlayments employed to retain packed aggregate, and components of composite roofing materials, insulation, pliable or flexible siding, and interior wall and ceiling finishes.

The highloft non-woven goods can be defined broadly as bonded, non-woven fibrous structures of varying bulks that provide varying degrees of resiliency, physical integrity, and durability depending on end use. Current major uses of highloft non-wovens include the manufacture of quilts, mattress pads, mattress covers, sleeping bags, furniture underlayments (padding), air filters, carpet underlayments (e.g., carpet pads), winter clothing, shoulder and bra pads, automotive, home and industrial insulation and paddings, padding and packaging for stored and shipped materials and otherwise hard surfaces (e.g., automobile roof tops, chairs, etc.), floor care pads for cleaning, polishing, buffing, and stripping, house robes (terrycloth, etc.), crib kick pads, furniture and toss pillows, molded packages, and kitchen and industrial scrub pads.

The polymers and methods can be used to manufacture all such non-wovens, and they are particularly useful for the manufacture of non-wovens free of, or having reduced levels of, formaldehyde or other potentially toxic components, and which have relatively high wet and dry tensile strength, abrasion resistance, color stability, stability to heat, light, detergent, and solvents, flexibility, elongation, shape retention, and/or acceptable "hand". The polymers are also particularly useful in manufacturing methods which require relatively short cure time (rapid bonding rate), relatively high polymer-to-fiber cohesion.

temperature stability (during curing and subsequent treatment), and/or the use of slightly acidic, neutral or alkaline application solutions or dispersions.

The method of the present invention can be used to manufacture articles that are suitable for use in applications, requiring sterilized products, e.g., the health care industry. Exemplary sterilized products include hospital gowns, masks, and bandages.

The polymers can be applied to the selected substrate by any of the procedures employed to apply other polymeric materials to the particular substrate. For example, a textile substrate can be immersed in the polymer solution or colloid in a typical dip-tank operation, sprayed with the polymer solution or colloid, or contacted with rollers or textile "printing" apparatus employed to apply polymeric colloids and solutions to textile substrates. Polymer concentration in the applied solution or colloid can vary considerably depending upon the procedures and apparatus employed to apply the polymer and the desired total polymer loading (polymer content) of the finished textile. Typically, the polymer concentration ranges from about 1 percent to about 60 percent. However, most applications involve solutions or colloids containing about 5 to about 60 weight percent latex solids.

Textile fiber assemblies wetted with substantial quantities of polymer solutions or latexes are typically squeezed with pad roll, knip roll, and/or doctor blade assemblies to remove excess solution or dispersion and, in some instances, to "break" and coalesce the latex and improve polymer dispersion, distribution, and fiber wetting.

The polymer is cross-linked by exposing the polymer/substrate combination to radiation capable of cross-linking the polymer. A rapid curing or cross-linking rate is important in essentially all methods of applying polymers to textiles and other substrates. For example, it is generally desirable to quickly reduce surface tackiness and increase fiber-to-fiber bond strength as soon as possible in the manufacture of loose woven textiles, knits, and non-wovens including all varieties of paper. Most often, adequate bond strength and sufficiently low surface tackiness must be achieved in textiles before they can be subjected to any significant stresses and/or subsequent processing. Accordingly, it is preferred that the polymer be capable of cross-linking when exposed to radiation for a period of less than about 15 minutes.

EXAMPLES

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In following examples, polymers containing pendant functional groups are synthesized. The polymers are applied to a substrate and cross-linked using UV light. The tensile strength of the UV cured polymer/substrate article is demonstrated.

Example 1

Preparation of Stock Polymer Latex

A surfactant-monomer pre-emulsion was formed by emulsifying about 5.3 g itaconic acid, about 10.6 g acrylamide, about 251.7 g butyl acrylate, about 255.8 g ethyl acrylate, about 32.7 g polyethoxylated nonylphenol surfactant containing about 40 moles ethylene oxide per mole, about 10.6 g polyethoxylated nonylphenol surfactant containing about 50 moles ethylene oxide per mole, and about 4.5 g sodium lauryl sulfate surfactant (about 30 percent active) in about 133.6 g water. A reactor was initially charged with about 353.4 g deionized water and about 1.1 g dissolved ammonium hydrogen phosphate to which about 70 ml of the monomer-surfactant pre-emulsion was then added. The resulting mixture was purged with nitrogen and heated to about 43° C. Sodium metabisulfite (about 0.45 g) and potassium peroxydisulfate (about 0.72 g) were then added with agitation, and the reactor was allowed to exotherm to about 60° C. The remainder of the monomer-surfactant pre-emulsion was then gradually metered into the reactor along with about 57 ml of a solution formed by dissolving about 4.8 g potassium peroxydisulfate in about 100 ml water and about 31 ml of a solution formed by dissolving about 4.4 g sodium metabisulfite in about 100ml water over a period of about 3 hours. The reactor's temperature was maintained at about 60 °C. throughout the reaction. Tertiary butyl hydroperoxide (about 0.4 g) was then added to assure polymerization of all monomers. The resulting latex contained about 48.4 weight percent latex solids and had a pH of about 2.9. The latex solids has a polymeric composition of about 1 weight percent itaconic acid, about 2 weight percent acrylamide, about 48 weight percent butyl acrylate, and about 49 weight percent ethyl acrylate based upon the total weight of the polymer.

Example 2

Preparation of Polymer Containing Pendant Functional Group

A latex of a polymer containing about 4 weight percent acetoacetoxyethylacrylate (AAEA) was prepared using the compositions and procedures described in Example 1, supra, with the exception that sufficient AAEA was incorporated in the monomer-surfactant pre-emulsion to form a polymer containing about 4 weight percent AAEA. The concentration of the other monomers was reduced proportionately to about 1 weight percent itaconic acid, about 1.9 weight percent acrylamide, about 46.1 weight percent butyl acrylate, and about 47 weight percent ethyl acrylate. All other compositions and conditions were as described in Example 1.

Example 3

Preparation of UV Cross-linked Polymer Coated Paper

Chromatographic grade filter paper was saturated with the polymer latex prepared in Example 2. The saturated filter paper was air dried. The dried filter paper was cut into 25 x 150 mm (one-inch by six-inch) strips. Each strip was vertically suspended approximately 175 mm (seven inches) away from eight circumferentially located, 460 mm (18-inch) fluorescent tubes. Adjacent fluorescent tubes were spaced approximately 45° apart. Each fluorescent tube was capable of emitting UV light having a wavelength of about 254 nm. As shown in Table I, infra, each strip of paper was exposed to either four equally spaced or all eight UV light sources for various times. While the latex polymer in each strip was being cured, the ambient temperature proximate the strip was as stated in Table I, infra.

Example 4

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Preparation of Heat Cured Polymer Coated Paper

Chromatographic grade filter paper was saturated with the polymer latex of Example 2. The saturated filter paper was air dried. The dried filter paper was cut in quarters. Each quarter section was heat cured at the temperatures stated in Table I, infra, for about 3 minutes. The oven dried sections were then cut into 25 x 150 mm (one-inch by six-inch) strips.

35 Example 5

Tensile Strength Test

Strips of the UV cross-linked sample of Example 3 or the heat cured strips of Example 4 were tested for wet tensile strength by dipping each strip in a 1 percent solution of Aerosol OT brand surfactant for about 4 seconds and measuring tensile on an Instron Model 1122 brand tensile gauge instrument. (Aerosol OT brand surfactant is manufactured by American Cyanamid, Inc.) Each wet tensile strength obtained is listed in Table I.

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TABLE I

| Cure Time, Minutes | WET TENSILE STRENGTH (kg (lb)) | | | | | |
|--------------------|--------------------------------|---------------------------|---------------------------|-------------------------------|--------------------------------|--|
| | Thermal Cure, 66 ° C. | Thermal Cure, 107 ° C. | Thermal Cure, 149 ° C. | UV Cure, 4/8 Tubes, 43 °C. | UV Cure, 8/8 Tubes, 51 ° C. | |
| 0 | 0.18(0.4) | 0.18(0.4) | 0.18(0.4) | 0.18(0.4) | 0.18(0.4) | |
| 0.5 | 0.45(1.0) | 0.54(1.2) | 1.09(2.4) | - | • | |
| 1 | - | <u>-</u> ` ` | - | 1.09(2.4) | 1.32(2.9) | |
| 3 | 0.45(1.0) | 0.82(1.8) | 2.27(5.0) | 1.27(2.8) | 1.41(3.1) | |
| 5 | 0.45(1.0) | 0.95(2.1) | 2.40(5.3) | _ | - | |
| 6 | 0.10(1.10) | . ` ` | - | 1.36(3.0) | 1.45(3.2) | |
| 10 | 0.45(1.0) | 1.27(2.8) | 2.72(6.0) | - | - | |
| 15 | 3.35(1.0) | | - ` ′ | 1.54(3.4) | 1.68(3.7) | |

The data of Table I demonstrate that UV light can effectively cure a polymer having pendant functional groups within the scope of formula I, supra. In addition, the data also indicate that the UV cured polymer improves the wet tensile strength of a treated substrate. Furthermore, a comparison of the wet tensile strength of UV cured articles with articles thermally cured at 66 °C. and 107 °C. indicates that UV cured articles attain a higher wet tensile strength in a shorter period of time than articles thermally cured at a much higher temperature. The ability of UV light to quickly cure articles at low temperatures helps avoid substrate degradation and makes UV curing a very attractive commercial process.

Example 6

Additional UV Cure Studies

Latexes of a polymer containing various amounts of AAEA were prepared using the compositions and procedures of Example 2, supra. Four separate portions of each latex were isolated and the pH of each portion was adjusted to either about 2, 4, 6 or 8. Strips of chromatographic grade filter paper were saturated with each pH-adjusted portion and cured according to the procedures of Examples 3 and 4. The wet tensile strengths of the cured polymer coated strips are set forth in Table II, infra.

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TABLE II (kg Wet Tensile Strength (lb) at 18% Add-On

| 5 | | | | | Heat | |
|----|-------|-----------------------|--|---|---------------------------|------|
| | | | UV Cure, | 8/8 Tubes, | Cured | |
| | | | at 44° | C., Min. | at 44°C. | |
| | AAEA, | Percent | | | For | Bath |
| | Wt. 8 | Add-On ¹ 0 | <u>0.5</u> <u>1</u> | 3 6 | 15 15 Min. | PH |
| 10 | 0 | 22.2 0.(8(0.4) | | 0.27 6.36 7 | 36 (0.8) 0.23 (0.5) | 2, |
| | 0 | 22.0 0.27(0.6) | | | (0.8)0.27(0.6) | 4 |
| 15 | 0 | 27.2 0.23(0.5) | | 27. 6. 6. 6. 6. | | 6 |
| | 0 | 26.0 6.23 (0.5) | ~~~.\$~\$~.\$ ~~~.\$~~\$ | 6.5,6.6 | (6.7°, 16.4) | 8 |
| 20 | 2 | 21.6 0.23 (0.5) | 0.5 | 332 0.41 0 | (1.6)0.23 (0.5) | 2 |
| 20 | 2 | 26.0 0.27 (0.6) | | | | 4 |
| | 2 | 24.1 0.32 (0.7) | | | | 6 |
| 25 | 2 | 22.0 0.32 (0.7) | \$. 8 . 8 . 8 . 8 . 8 . 8 . 8 . 8 . 8 . | 1. 4. 0. E. 19. | (1.4)° (0.9) | 8 |
| | 5 | | 5.23 6.23 ((0.5) 6.5) | | (1.5) (0.5) | 2 · |
| 30 | 5 | 25.6 0.27 (0.6) | 0.22 (0.6) (0.7) | (1.0) (1.3) | (2.8) (2.7) | 4 |
| | 5 | 21.7 0.8 | 3.41 .9 (1.0) | (1.3), (1.5), | (2.0) (6.9) | 6 |
| | 5 | 24.8 0.32 0.7 | 3.36.80.36.8 | (t. 0, t. 0, | (E. 9) (E. 7) | 8 |
| 35 | 10 | 24.8 0.18(0.4) | | 1.32 0.55 | (1.8) (0.4) | 2 |
| | 10 . | 23.3 0.32 (0.7) | 3 3 7 (0. 8) (0. 8) | 1. 2. 4. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. | (1.6) (0.7) | 4 |
| 40 | 10 | 27.3 0.37 | 56. P. 82.7 | | (1.3) (0.7) | 6 |
| | 10 | 25.7 0.22 (0.7) | 5.36.8°,36.8 | 7(1·0) (1·3) | (1.5)0.5 (0.8) | 8 |

Percent add-on can be represented by the formula (Y - X)/X 100% wherein X is the initial weight of a strip of chromatographic filter paper and Y is the weight of the cured, polymer coated filter paper.

The data set forth in Table II demonstrate that polymers prepared without AAEA, an exemplary, monomer within the scope of formula VIII, supra, failed to significantly improve tensile strength as a result of UV curing. The data also sustantiate that UV cured articles attain a higher wet tensile strength in a shorter period of time than articles thermally cured at the same temperature.

Example 7

Further UV Cure Studies

Cured polymer coated filter paper strips were prepared as described in Example 6 with one modification. The modification comprised forming the surfactant-monomer pre-emulsion by emulsifying about 4.00 g itaconic acid, about 7.95 g acrylamide, about 188.78 g butyl acrylate, about 191.85 g ethyl acrylate, and about 130.85 g vinyl acetate. The non-monomer ingredients of the surfactant-monomer pre-emulsion were the same as employed in Example 1. The wet tensile strengths of cured polymer coated strips prepared in accordance with this example are set forth in Table III, infra.

Wet Tensile Strength (1b) at 18% Add-On

| | | | UV Cure, 8/8 Tubes, at 44° C., Min | Heat Cured at 44°C. |
|----|---------------|---------------------|--|----------------------------|
| 10 | AAEA, Wt.% | Percent Add-On 0 | 0.5 1 3 6 15 | For Bath 15 Min. pH |
| 10 | 0 | 20.2 0.5 (1.3) | | 2 (6.3) |
| | 0 | | 1.3°.27.2°,27.2°,27.3°,44.4 | 0.54 |
| 15 | 0 | 20.3 0,57 (1.2) | | 0,34(1.3) 6 |
| | 0 | 20.0015 (1.3) | 1.3 (1.3) (1.3) (1.4) (1.4) | 0.5 1 10.2 8 |
| 20 | 2 | 20.5 0.5 (1.3) | .57 0.59 0.64 0.68 0.91 (1.2) (1.3) (1.4) (1.5) (2.0 | 0.54 |
| | 2 | 20.6 0.5 (1.2) | A. 3. 8. 5. 6. 5. 6. 6. 6. 6. 6. 6. 6. 6. 6. 6. 6. 6. 6. | 0,59 |
| | 2 | | 13. 6° 7. 6° 17. 7° 8° 8° 12. 1 | |
| 25 | 2 | 19.9 | 6. 5, (1. 6, 7), (1. 8, 2. 1) | 016(1.4) 8 |
| | 5 | 17.5 0.68 (1.5) | 1.5) 1.5) 1.6) 1.5A | 0.64.4) 2 |
| 30 | 5 | 17.5 0.68 (1.5) | 13.6 1. 6 1. 6 1. 6 2. 1 2. 6 2. 1 2. 6 2. 6 | 0.65(1.5) 4 |
| | 5 | | 1,13 0, 15 0,26 0,41 0,18.6 | |
| 35 | 5 | 18.1 | 1. 9, 1. 8, 1. 8, 2. 9, 2. 4 | 0.64 8 |
| | 10 | 18.1 (1.3) | 1.30 (1.3) (1.5) (1.9) (2.0) | 0.59 |
| 40 | 10 | | (6) 0,60 0,77 0,65 1,27 (1. 7) (2.1) (2.8) | |
| | 10 | | 17 . 7 . 17 . 7 . 8 . 6 . 9 . 0 . 12 . 6 . 6 . 6 . 6 . 6 . 6 . 6 . 6 . 6 . | 0.72 6 |
| 45 | 10 | 17.5 0.87 (1.8) | 1.7, 1.7, 2.9, 2.2, 2.8 | (1.6) 8 |

See Table II, footnote 1, supra.

The data set forth in Table III support the observations made regarding the data of Table II, supra.

Although the present invention has been described in considerable detail with reference to certain preferred versions thereof, other versions are possible. For example, the polymers can be coated on substrates other than those specifically mentioned above. In addition, the polymer need not be coated onto a substrate in order to be UV cured. Instead, the polymer can be confined to a desired shape and then UV cured.

Claims

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1. A method of producing an article comprising shaping a polymer comprising a backbone and a plurality of pendant functional groups having the formula:

wherein R_1 is a divalent radical and X is organoacyl or cyano; and curing the shaped polymer by exposure to radiation having a wavelength within the range of about 10^{-3} to about 265 nm.

2. A method of producing an article comprising
(a) contacting a substrate with a polymer to form a substrate/polymer combination, the polymer comprising a backbone and a plurality of pendant functional groups, the pendant functional groups having the formula:

wherein R_1 is a divalent radical and X is organoacyl or cyano; and (b) curing the polymer by exposing the substrate/polymer combination to radiation having a wavelength within the range of about 10^{-3} to about 265 nm.

- 3. A method according to claim 1 or 2, wherein the radiation is ultraviolet light.
- 4. A method according to claim 3, wherein the ultraviolet light has a wavelength in the range of about 10 to about 265 nm.
 - 5. A method according to claim 4, wherein the ultraviolet light has a wavelength not more than about 250 nm.
 - A method according to claim 5, wherein the ultraviolet light has a wavelength of not more than about 225 nm.
- A method according to claim 1 or 2, wherein the radiation is ultraviolet light having a wavelength of
 about 225 to about 265 nm.
 - 8. A method according to any preceding claim, wherein the shaped polymer or the substrate/polymer combination is exposed to the radiation for a period of time of up to about 15 minutes.
- 45 9. A method according to any preceding claim, wherein the polymer is a constituent of a solution.
 - 10. A method according to any one of claims 1 to 8, wherein the polymer is a constituent of a colloid.
 - 11. A method according to claim 10, wherein the colloid has a pH of about 2 to about 8.
 - 12. A method according to any preceding claim, wherein pendant functional groups have the formula

wherein

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X is organoacyl;

Y is selected from oxygen, sulfur and NR4;

Z is selected from oxygen, sulfur and NR4;

R₃ is a divalent organic radical at least 2 atoms in length; and

Re is selected from hydrogen and hydrocarbyl radicals having up to 6 carbon atoms.

- 13. A method according to claim 12, wherein R₃ comprises up to about 40 atoms.
- 10 14. A method according to claim 13, wherein R₃ comprises up to about 20 atoms.
 - 15. A method according to claim 12, wherein R₃ is selected from substituted and unsubstituted alkylenes, polyoxyalkylenes, polythiolalkylenes and polyaminoalkylenes.
 - 16. A method according to claim 12, wherein R₃ is selected from

$$- (R_{6})_{m} \xrightarrow{- (R_{7})_{n} - ,}$$

$$- (R_{6})_{m} \xrightarrow{- (R_{2}CH_{3})_{n} - ,} \text{ and}$$

$$- (R_{6})_{m} - \frac{CH_{2}CH_{3}}{CH_{2}CH_{3}} (R_{7})_{n} - ,$$

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wherein

m is 0 or 1;

n is 0 or 1;

 R_δ is a divalent organic radical at least one atom in length; and R_7 is a divalent organic radical at least one atom in length.

17. A method according to any preceding claim, wherein X is

- 45 R₂ being selected from hydrogen and monovalent organic radicals.
 - 18. A method according to claim 17, wherein R₂ contains up to about 10 atoms in addition to any hydrogen atoms present in the monovalent organic radical.
- 50 19. A method according to claim 17, wherein R₂ is methyl.
 - 20. A method according to any preceding claim, further comprising the step of forming the backbone by combining one or more olefinically unsaturated monomers.
- 55 21. A method according to claim 20, wherein at least one of the olefinically unsaturated monomers has the formula

$$R_{Q} - CH = C - R_{1} - C - CH_{2} - X$$

wherein $R_{\rm 9}$ and $R_{\rm 10}$ are each independently selected from hydrogen, halo, thio, and monovalent organic radicals.

22. A method according to claim 21, wherein X is

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and R₁ is a divalent organic radical of at least 3 atoms in length.

23. A method according to claim 22, wherein the polymerizable functional monomer has the formula selected from

$$CH_{2} = C - C - 0 - CH_{2}CH_{2} - 0 - C - CH_{2} - C - CH_{3},$$

$$CH_{2} = \overset{C}{C} - \overset{O}{C} - O - CH_{2}CH_{2}CH_{2} - O - \overset{O}{C} - CH_{2} - \overset{O}{C} - CH_{3}$$
, and

$$CH_{3} = \overset{CH_{3}}{\overset{0}{\text{ll}}} - \overset{O}{\overset{C}{\text{ll}}} - CH_{2} - \overset{CH_{2}CH_{3}}{\overset{C}{\text{ll}}} - O - \overset{O}{\overset{C}{\text{ll}}} - CH_{2} - \overset{O}{\overset{C}{\text{ll}}} - CH_{3}$$

- 24. A method according to claim 21 or 22, wherein at least one of the monomers is selected from (a) monocarboxylic acid esters, (b) polycarboxylic acid esters, (c) salts of (a) and (b), and (d) mixtures thereof.
 - 25. A method according to claim 21, wherein at least one of the monomers is selected from acrylic acid, itaconic acid, hydroxyethyl acrylate, acrylonitrile, and acrylamide.
 - 26. A method according to any preceding claim, wherein the polymer comprises at least about 0.5 weight percent of the pendant functional groups.
- 27. A method according to claim 24, wherein the polymer comprises up to about 20 weight percent of the pendant functional groups.
 - 28. A method according to claim 2 or any claim dependent thereon, wherein the substrate is a textile.

Patentansprüche

 Verfahren zur Herstellung eines Gegenstandes, dadurch gekennzeichnet, daß man ein Polymer formt, das ein Rückgrat und eine Mehrzahl von daran hängenden funktionellen Gruppen der Formel

$$-R_1 - C - CH_2X$$

aufweist, worin R₁ ein zweiwertiger Rest ist und X ein Organoacyl oder Cyano ist und das geformte Polymere härtet, indem man es einer Strahlung mit einer Wellenlänge im Bereich von etwa 10⁻³ bis etwa 265 nm aussetzt.

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2. Verfahren zur Herstellung eines Gegenstandes, dadurch gekennzeichnet, daß man (a) eine Unterlage mit einem Polymeren in Kontakt bringt, um eine Kombination Unterlage/Polymer zu bilden, wobei das Polymere ein Rückgrat und eine Mehrzahl von daran hängenden funktionellen Gruppen aufweist, wobei die daran hängenden funktionellen Gruppen die Formel

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aufweisen, worin R1 ein zweiwertiger Rest ist und X ein Organoacyl oder Cyano ist, und (b) das Polymere härtet indem man die Kombination Unterlage/Polymeres einer Strahlung mit einer Wellenlänge im Bereich von etwa 10⁻³ bis etwa 265 nm aussetzt.

Verfahren nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß die Strahlung ultraviolettes Licht ist.

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Verfahren nach Anspruch 3, dadurch gekennzeichnet, daß das ultraviolette Licht eine Wellenlänge im Bereich von etwa 10 bis etwa 265 nm hat.

Verfahren nach Anspruch 4, dadurch gekennzeichnet, daß das ultraviolette Licht eine Wellenlänge von nicht mehr als etwa 250 nm hat.

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6. Verfahren nach Anspruch 5, dadurch gekennzeichnet, daß das ultraviolette Licht eine Wellenlänge von nicht mehr als etwa 225 nm hat.

7. Verfahren nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß die Strahlung ultraviolettes Licht mit einer Wellenlänge von etwa 225 bis etwa 265 nm ist.

8. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß das geformte Polymere oder die Kombination Unterlage/Polymeres der Strahlung für eine Zeitspanne von bis zu etwa 15 Minuten ausgesetzt wird.

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9. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß das Polymere ein Bestandteil einer Lösung ist.

10. Verfahren nach einem der Ansprüche 1 bis 8, dadurch gekennzeichnet, daß das Polymere ein Bestandteil eines Kolloids ist.

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11. Verfahren nach Anspruch 10, dadurch gekennzeichnet, daß das Kolloid ein pH von etwa 2 bis etwa 8

12. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß daran hängende funktionelle Gruppen die Formel

$$-\overset{0}{c} - Y - R_3 - Z - \overset{0}{c} - CH_2X$$

haben, worin

X ein Organoacyl;

Y Sauerstoff, Schwefel oder NR4;

Z Sauerstoff, Schwefel oder NR4;

R₃ ein zweiwertiger organischer Rest von wenigstens 2 Atomen Länge und

Ru Wasserstoff oder Hydrocarbylreste mit bis zu 6 Kohlenstoffatomen sind.

- 13. Verfahren nach Anspruch 12, dadurch gekennzeichnet, daß R₃ bis zu etwa 40 Atome aufweist.
- 10. Verfahren nach Anspruch 13, dadurch gekennzeichnet, daß R₃ bis zu etwa 20 Atome aufweist.
 - 15. Verfahren nach Anspruch 12, dadurch gekennzeichnet, daß R₃ aus der Gruppe substituierte und unsubstituierte Alkylene, Polyoxyalkylene, Polythioalkylene und Polyaminoalkylene ausgewählt ist.
- 15 16. Verfahren nach Anspruch 12, dadurch gekennzeichnet, daß R3 aus den folgenden Formein gewählt ist

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worin

m die Zahl 0 oder 1;

n die Zahl 0 oder 1:

Rs ein zweiwertiger organischer Rest mit einer Länge von wenigstens einem Atom und

R₇ ein zweiwertiger organischer Rest mit einer Länge von wenigstens einem Atom sind.

17. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß X die Gruppe

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ist, worin R2 Wasserstoff oder ein einwertiger organischer Rest ist.

- 18. Verfahren nach Anspruch 17, dadurch gekennzeichnet, daß R₂ bis zu etwa 10 Atome zusätzlich zu allen im einwertigen organischen Rest vorhandenen Wasserstoffatomen enthält.
- 19. Verfahren nach Anspruch 17, dadurch gekennzeichnet, daß R2 Methyl ist.
- Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß es weiterhin die Stufe der Bildung des Rückgrats durch Kombination von einem oder mehreren olefinisch ungesättigten Monomeren umfaßt.
- 21. Verfahren nach Anspruch 20, dadurch gekennzeichnet, daß wenigstens eines der olefinisch ungesättigten Monomeren die Formel

$$R_{9} - CH = C - R_{1} - C - CH_{2} - X$$

hat, worin

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 R_9 und R_{10} jeweils unabhängig voneinander Wasserstoff, Halogen, Thio oder einwertige organische Reste sind.

22. Verfahren nach Anspruch 21, dadurch gekennzeichnet, daß X der Rest

ist und R₁ ein zweiwertiger organischer Rest mit einer Länge von wenigstens 3 Atomen ist.

23. Verfahren nach Anspruch 22, dadurch gekennzeichnet, daß das polymerisierbare funktionelle Monomere eine der folgenden Formeln aufweist:

$$_{\text{ch}}^{\text{CH}} = \text{C} - \text{C} - \text{O} - \text{CH}_{2}\text{CH}_{2} - \text{O} - \text{C} - \text{CH}_{2} - \text{C} - \text{CH}_{3}$$

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 CH₂ = $^{\text{CH}}_{2}$ $^{\text{CH}}_{2}$ $^{\text{CH}}_{2}$ CH₂ - $^{\text{CH}}_{2}$ - $^{\text{CH}}_{2}$ - $^{\text{CH}}_{2}$ - $^{\text{CH}}_{2}$ $^{\text{Oder}}$

- 24. Verfahren nach Anspruch 21 oder 22, dadurch gekennzeichnet, daß wenigstens eines der Monomeren aus (a) Monocarbonsäureestern, (b) Polycarbonsäureestern, (c) Salzen von (a) und (b), und (d) Gemischen davon ausgewählt ist.
- 25. Verfahren nach Anspruch 21, dadurch gekennzeichnet, daß wenigstens eines der Monomeren aus der Gruppe Acrylsäure, Itaconsäure, Hydroxyethylacrylat, Acrylnitril und Acrylamid ausgewählt ist.
 - 26. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß das Polymere wenigstens etwa 0.5 Gew.-% der daran hängenden funktionellen Gruppen enthält.
 - 27. Verfahren nach Anspruch 24, dadurch gekennzeichnet, daß das Polymere bis zu etwa 20 Gew.% der daran hängenden funktionellen Gruppen enthält.
- 28. Verfahren nach Anspruch 2 oder einem davon abhängigen Anspruch, dadurch gekennzeichnet, daß die
 Unterlage ein Textilmaterial ist.

Revendications

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1. Une méthode pour produire un article comprenant : la conformation d'un polymère comportant un squelette et plusieurs groupes fonctionnels pendants ayant la formule :

dans laquelle R_1 est un radical divalent et X est organoacyl ou cyano ; et le durcissement du polymère conformé en l'exposant à un rayonnement ayant une longueur d'onde comprise dans la gamme d'environ 10^{-3} à environ 265 nm.

2. Une méthode pour produire un article comprenant : (a) la mise en contact d'un substrat avec un polymère pour former une combinaison substrat/polymère, le polymère comprenant un squelette et plusieurs groupes fonctionnels pendants , les groupes fonctionnels pendants ayant la formule:

dans laquelle R₁ est un radical divalent et X est organoacyl ou cyano; et

(b) le durcissement du polymère en exposant la combinaison substrat/polymère à un rayonnement ayant une longueur d'onde compris dans la gamme d'environ 10⁻³ à environ 265 nm.

- 3. Une méthode selon la revendication 1 ou 2, dans laquelle le rayonnement est de la lumière ultraviolette
- Une méthode selon la revendication 3 , dans laquelle la lumière ultraviolette a une longueur d'onde dans la gamme d'environ 10 à environ 265 nm.
- Une méthode selon la revendication 4, dans laquelle la lumière ultraviolette a une longueur d'onde ne dépassant pas environ 250 nm.
 - Une méthode selon la revendication 5 , dans laquelle la lumière ultraviolette a une longueur d'onde ne dépassant pas environ 225 nm.
- 7. Une méthode selon la revendication 1 ou 2, dans laquelle le rayonnement est de la lumière ultraviolette ayant une longueur d'onde d'environ 225 à environ 265 nm.
 - 8. Une méthode selon l'une quelconque des revendications précédentes, dans laquelle le polymère conformé ou la combinaison substrat/polymère est exposé au rayonnement pendant une période de temps allant jusqu'à environ 15 minutes.
 - 9. Une méthode selon l'une quelconque des revendications précédentes, dans laquelle le polymère est le constituant d'une solution .
- 50 10. Une méthode selon l'une quelconque des revendications 1 à 8 , dans laquelle le polymère est le constituant d'un colloïde .
 - 11. Une méthode selon la revendication 10, dans laquelle le colloïde a un pH d'environ 2 à environ 8.
- 55 12. Une méthode selon l'une quelconque des revendications précédentes , dans laquelle les groupes fonctionnels pendants ont la formule :

dans laquelle :

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X est organoacyl;

Y est choisi parmi oxygène, soufre et NR4;

Z est choisi parmi oxygène, soufre et NR4;

R₃ est un radical organique divalent ayant au moins la longueur de 2 atomes ; et

R4 est choisi parmi hydrogène et les radicaux hydrocarbyl ayant jusqu'à 6 atomes carbone .

- 13. Une méthode selon la revendication 12 , dans laquelle $R_{\rm 3}$ comprend jusqu'à 40 atomes .
- 5 14. Une méthode selon la revendication 13, dans laquelle R₃ comprend jusqu'à 20 atomes.
 - 15. Une méthode selon la revendication 12, dans laquelle R₃ est choisi parmi les alcoylènes , polyoxyalcoylènes, polythiolalcoylènes et polyaminoalcoylènes, substitués et non substitués.
- 16. Une méthode selon la revendication 12, dans laquelle R₂ est choisi parmi :

$$-(R_6)_m$$
 $(R_7)_n$ $-$

dans lesquels:

m est O ou 1;

n est O ou 1:

 R_{ϵ} est un radical organique divalent ayant au moins la longueur d'un atome ; et

 $\ensuremath{\mathsf{R}}_7$ est un radical organique divalent ayant au moins la longueur d'un atome .

17. Une méthode selon l'une quelconque des revendications précédentes, dans laquelle X est

- 50 R₂ étant choisi parmi hydrogène et les radicaux organiques monovalents.
 - 18. Une méthode selon la revendication 17, dans laquelle R₂ contient jusqu'à 10 atomes en plus de tout atome d'hydrogène présent dans le radical organique monovalent.
- 55 19. Une méthode selon la revendication 17, dans laquelle R_2 est méthyl .
 - 20. Une méthode selon l'une quelconque des revendications précédentes, comprenant en outre la phrase consistant à former le squelette en combinant un ou plusieurs monomères oléfiniquement non-saturés.

21. Une méthode selon la revendication 20 , dans laquelle au moins un des monomères oléfiniquement non-saturés a la formule :

$$R_9 - CH = C - R_1 - C - CH_2 - X$$

dans laquelle R₉ et R₁₀ sont chacun, indépendamment, choisis parmi hydrogène , halo, thio et les radicaux organiques monovalents .

22. Une méthode selon la revendication 21 , dans laquelle X est

et R_1 est un radical organique divalent ayant la longueur d'au moins 3 atomes .

23. Une méthode selon la revendication 22, dans laquelle le monomère fonctionnel polymérisable a une formule choisie parmi :

$$_{\text{CH}_2}^{\text{CH}_3}$$
 $_{\text{C}}^{\text{O}}$ $_{\text{C}}^{\text{CH}_2}$ $_{\text{C}}^{\text{C}}$ $_{\text{C}}^{\text{C$

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- 24. Une méthode selon la revendication 21 ou 22, dans laquelle au moins un des monomères est choisi parmi (a) les esters d'acides monocarboxyliques, (b) les esters d'acides polycarboxyliques, (c) les sels de (a) et (b), et (d) les mélanges de ceux-ci.
- 25. Une méthode selon la revendication 21, dans laquelle au moins un des monomères est choisi parmi acide acrylique, acide itaconique, acrylate d'hydroxyéthyle, acrylonitrile et acrylamide.
- 26. Une méthode selon l'une quelconque des revendications précédentes, dans laquelle le polymère comprend au moins environ 0,5 % en poids de groupes fonctionnels pendants.
 - 27. Une méthode selon la revendication 24, dans taquelle le polymère comprend jusqu'à 20 % en poids de groupes fonctionnels pendants .
- 28. Une méthode selon la revendication 2 ou toute revendication dépendant de celle-ci, dans laquelle le substrat est un textile.